



Looking inside a Ni-Fe/MgAl₂O₄ catalyst for methane dry reforming via Mössbauer spectroscopy and in situ QXAS

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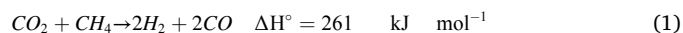
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ABSTRACT

The evolution of the constituents of an 8 wt%Ni-5 wt%Fe/MgAl₂O₄ catalyst for dry reforming of methane (DRM) is monitored by in situ quick X-ray absorption spectroscopy (QXAS) and ⁵⁷Fe Mössbauer spectroscopy. In as prepared state, Fe is present as NiFe₂O₄ at the surface and as MgFe³⁺_xAl_{2-x}O₄ within the support, whereas Ni is mainly present as NiO. During H₂-TPR, NiFe₂O₄ and NiO form an alloy from 500 °C on and MgFe³⁺_xAl_{2-x}O₄ is partially reduced to MgFe²⁺_xAl_{2-x}O₄, such that Ni-Fe alloy, MgFe²⁺_xAl_{2-x}O₄ and MgFe³⁺_xAl_{2-x}O₄ are the prevalent phases in the reduced catalyst. During DRM, dominantly oxidizing environments (CH₄/CO₂ = 1/2, 1/1.5) lead to formation of FeO_x nanoparticles at the surface of the Ni-Fe alloy, thereby affecting the DRM activity, and possibly to some reincorporation of Fe into the support. For CH₄/CO₂ = 1/1, no significant changes occur in the catalyst's activated state, as a consequence of reduction by CH₄ dissociation species counteracting oxidation by CO₂. However, Mössbauer analysis detects continued extraction of Fe from the support, sustaining ongoing Ni-Fe alloying.

1. Introduction

DRM, represented in Eq. (1), is of major interest for sustainable development as it entails the catalytic conversion of the greenhouse gases CO₂ and CH₄ to CO and H₂, i.e. syngas, which can serve as building blocks in the creation of high-value end products, e.g. via Fischer Tropsch [1,2] or methanol synthesis processes [3,4].



As global supply of natural gas has increased tremendously [5], methane has become an economically interesting co-reactant for CO₂ conversion, making DRM an even more attractive pathway for CO₂ utilization. DRM has a low operating cost compared to other methane reforming processes such as steam reforming [6,7], autothermal reforming [8,9] and partial oxidation of methane [10,11].

Despite its advantages, different obstacles impede the industrial exploitation of DRM. The high endothermicity of Eq. (1) requires the use

of high temperatures (650–850 °C). With equimolar CH₄ and CO₂ concentrations, i.e. industrially relevant feed composition, this promotes catalyst deactivation due to coke formation and sintering of the support and metal particles [12–14]. It is therefore mandatory that an efficient DRM catalyst is thermally stable and resistant to coke formation and sintering, whilst also being highly active and cost-effective.

Ni-Fe catalysts have proved successful in meeting the above criteria, leading to a rise in the application of these materials in reforming reactions. Wang et al. [15] studied Ni-Fe/Al₂O₃ for steam reforming of biomass-derived tar, and concluded that the catalyst displayed an enhanced performance over monometallic Ni and Fe counterparts. This was attributed to the formation of a Ni-Fe alloy, wherein synergy between Ni as tar activation site and Fe as oxygen carrier that oxidizes the carbon species formed on neighboring Ni sites decreases coke accumulation. Analogous Ni-Fe synergistic effects have been reported in methanation [16], chemical looping [17] and furfural conversion [18] catalysts. Kim and co-workers [19] compared monometallic Ni, Fe and

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bimetallic Ni-Fe catalysts on a hydrotalcite-derived $\text{Mg}_x\text{Al}_y\text{O}_z$ support for DRM at 650 °C, and found that catalysts with $\text{Ni}/(\text{Fe}+\text{Ni}) = 0.8$ displayed the highest activity and stability. Ex situ and in situ characterization showed the redox functionality of the Fe species suppressed carbon formation in DRM. These results are in accordance with the work of Theofanidis et al. [20], who attributed the coke resistance in Ni-Fe/ MgAl_2O_4 DRM catalysts to the formation of FeO_x at the alloy surface, which react via a redox mechanism with the deposited carbon.

In spite of these studies, questions remain as to the nature of the phases present in these Ni-Fe reforming catalysts after reduction/activation and during reaction. Using operando XAS, Kim and co-workers [19] observed the formation of FeO in hydrotalcite-derived Ni-Fe/ $\text{Mg}_x\text{Al}_y\text{O}_z$ during DRM at 650 °C and 1 atm and equimolar CH_4 and CO_2 concentrations. For spinel-supported Ni-Fe/ MgAl_2O_4 , the formation of FeO_x has only been confirmed via in situ X-ray diffraction (XRD) during DRM (at 750 °C and 1 atm) at conditions with relatively high CO_2 partial pressures, i.e. $\text{CH}_4/\text{CO}_2 \leq 1/6$ [20]. At feed compositions closer to equimolar CH_4 and CO_2 concentrations, which are of high industrial importance, these oxides cannot be detected by diffraction. Hence, it is unclear whether this is because (i) such oxide phase indeed does not form in Ni-Fe/ MgAl_2O_4 under such conditions, (ii) the corresponding crystallites are too small for detection via XRD, or (iii) their concentrations are too low for detection via this technique. The presence of FeO is of particular importance as the latter was found to be at the origin of enhanced sintering given its low Tamman temperature [21].

Another matter worthy of investigation is the Ni-Fe alloy formation process during catalyst activation. Due to the overlapping nature of characteristic diffraction peaks, the presence of NiO, $\gamma\text{-Fe}_2\text{O}_3$, or NiFe_2O_4 cannot be confirmed beyond doubt by XRD studies [20]. In addition, dominant contributions of MgAl_2O_4 prevent detection of a possible NiAl_2O_4 phase. Moreover, spinel supports like MgAl_2O_4 can incorporate Fe, typically following one pot co-precipitation synthesis [22–24], yielding contributions related to Fe-substituted MgAl_2O_4 phases. For the present Ni-Fe/ MgAl_2O_4 impregnated catalysts however, such Fe-incorporated support phase lacks confirmation. Resolving these uncertainties will allow a better understanding of the behavior of these catalysts under DRM reaction conditions and lay the path towards improved catalyst synthesis.

In the present work, the above structural questions are addressed by means of a QXAS study of a MgAl_2O_4 -supported DRM catalyst with 8 wt % Ni and 5 wt% Fe, hereafter called 8Ni5Fe/ MgAl_2O_4 . This Ni/Fe ratio was chosen as it shows optimal activity and stability, with limited sintering and low carbon formation [20]. To characterize the sample during reaction, in situ QXAS [25] is employed at the Ni K and Fe K edge, both for H_2 -TPR and DRM under industrially relevant conditions, i.e. $\text{CH}_4/\text{CO}_2 = 1/1$, 750 °C and $\text{GHSV} = 4580 \text{ h}^{-1}$. Speciation of time-resolved XAS data is performed by means of statistical analysis via principal component analysis (PCA) and multivariate curve resolution coupled with alternating least squares (MCR-ALS). The analysis of the QXAS data is complemented with both transmission and surface-sensitive ^{57}Fe Mössbauer spectroscopy in as prepared state, i.e. after calcination, as well as after H_2 -TPR and after DRM.

2. Materials and methods

2.1. Catalyst preparation

MgAl_2O_4 support was produced via coprecipitation of Al ($(\text{NO}_3)_3 \cdot 9 \text{H}_2\text{O}$ (ACS reagent, $\geq 98\%$, Sigma-Aldrich) and Mg ($(\text{NO}_3)_2 \cdot 6 \text{H}_2\text{O}$ ($\geq 99\%$, Sigma-Aldrich) as described in the work of Theofanidis et al. [20]. To acquire metal loadings of 8 wt% Ni and 5 wt% Fe, appropriate amounts of the corresponding nitrates were used in an incipient wetness impregnation method. Details on the catalyst preparation and preliminary characterization of composition have been reported previously [20].

2.2. XAS references preparation

Fe^{3+} -substituted MgAl_2O_4 ($\text{MgFe}^{3+}\text{AlO}_4$) with 10 wt% Fe content was prepared via coprecipitation of $\text{Al}(\text{NO}_3)_3 \cdot 9 \text{H}_2\text{O}$ (ACS reagent, $\geq 98\%$, Sigma-Aldrich), $\text{Mg}(\text{NO}_3)_2 \cdot 6 \text{H}_2\text{O}$ ($\geq 99\%$, Sigma-Aldrich) and Fe ($(\text{NO}_3)_3 \cdot 9 \text{H}_2\text{O}$ ($\geq 99.95\%$, Sigma-Aldrich) as reported by Dharanipragada et al. [23]. NiFe_2O_4 and NiAl_2O_4 references were synthesized through a sol-gel autocombustion method. In a typical synthesis, 0.551 g $\text{Ni}(\text{NO}_3)_2 \cdot 6 \text{H}_2\text{O}$ (99.999%, Sigma-Aldrich) (1.90 mmol) was put to a 100 mL glass beaker containing 7 mL deionized water. Then, the appropriate quantity of $\text{Al}(\text{NO}_3)_3 \cdot 9 \text{H}_2\text{O}$ (ACS reagent, $\geq 98\%$, Sigma-Aldrich) or $\text{Fe}(\text{NO}_3)_3 \cdot 9 \text{H}_2\text{O}$ ($\geq 99.95\%$, Sigma-Aldrich) was added to obtain a molar ratio of Ni/Al (or Ni/Fe) of 1/2, as well as citric acid (ACS reagent, $\geq 99.5\%$, Sigma-Aldrich), such that $n_{\text{citric acid}}/n_{\text{nitrates}} = 1/1$. The aqueous solution was magnetically stirred at 50 °C at 350 rpm, and 50 mg of polyvinylpyrrolidone (PVP; average mol weight 40,000, Sigma-Aldrich) was added. The mixture was left to stir at constant temperature until all PVP was dissolved, after which the temperature was increased to 100 °C and maintained until a transparent gel was formed. This gel was dried overnight at 120 °C, yielding a powder that was first calcined under air flow at 400 °C (3 °C/min) for 1 h, and further up to 1000 °C (3 °C/min) with a 1 h dwell time.

2.3. Characterization techniques

2.3.1. QXAS measurements

In situ transmission QXAS measurements were performed at the ROCK beamline of the French synchrotron SOLEIL [26]. The storage ring was operated at 2.75 GeV with a ring current of 500 mA in top-up mode. A Frahm monochromator [25] equipped with a Si(111) channel-cut crystal oscillated at 2 Hz and scanned both the Fe K (7112 eV) and Ni K (8333 eV) edges, using a macro for fast edge switching. This resulted in quasi-simultaneous acquisition of both edges, with a delay of ~ 60 s. Catalytic samples of approximately 5 mg, ground to powders and 50% diluted with boron nitride, were inserted into a 2 mm quartz capillary reactor in between two quartz wool plugs. The capillary reactor was implemented in a frame which was connected to gas feed lines through Swagelok fittings. The inlet gas flow rates were maintained by means of calibrated Brooks mass flow controllers and a total flow between 10 and 60 NmL/min was employed. An external, calibrated heat gun was used to reach reaction temperatures up to 800 °C. XAS spectra energy calibration was assured by the measurement of Ni and Fe foils.

H_2 -TPR experiments were performed by subjecting the sample to a heating rate of 10 °C/min from 20 °C to a maximum of 800 °C, with a holding time of 30 min, under a flow of 10 NmL/min of 5 V% H_2 in He. After this reduction program, an in situ DRM measurement was performed with a molar CH_4/CO_2 ratio of 1/2 (2.6 NmL He/min, 4.2 NmL CH_4 /min and 8.4 NmL CO_2 /min) at 750 °C and 1 atm for 30 min. Following this DRM measurement, the catalyst was regenerated, i.e. brought back into its metallic state, by an isothermal H_2 reduction (5 V% H_2 in He, 15 NmL/min) at 750 °C for 30 min, after which it was subjected to a second DRM measurement with a molar CH_4/CO_2 ratio of 1/1.5 (3.0 NmL He/min, 4.6 NmL CH_4 /min and 6.8 NmL CO_2 /min) at 750 °C and 1 atm for 30 min. Following regeneration by an isothermal H_2 treatment, a final in situ DRM measurement was carried out for a molar CH_4/CO_2 ratio of 1/1 (3.0 NmL He/min, 5.0 NmL CH_4 /min and 5.0 NmL CO_2 /min), also at 750 °C and 1 atm for 30 min. For ease of expression, the identity $R_c = \text{CH}_4/\text{CO}_2$ will be used hereafter.

The same setup was used for ex situ measurements of reference samples. Oxidized state references for Fe and Ni were obtained via measurements of pelletized $\alpha\text{-Fe}_2\text{O}_3$, $\gamma\text{-Fe}_2\text{O}_3$, NiFe_2O_4 , $\text{MgFe}^{3+}\text{AlO}_4$, Fe_3O_4 , FeO, NiAl_2O_4 and NiO.

20 consecutive scans of the in situ QXAS data were averaged to increase the signal to noise ratio. All recorded data were normalized via the Python normalization GUI created by O. Roudenko [27]. Linear combination fitting (LCF) of the normalized XANES spectra of the as

prepared catalyst was performed using the associated module in the Athena software [28]. Analysis was restricted to the energy regions [7111 – 7161] eV and [8327 – 8377] eV for Fe K and Ni K edge data, respectively.

For the purpose of speciating the different Ni and Fe species through MCR-ALS analysis of the QXAS datasets, the number of principal components in each QXAS dataset was assessed through PCA [29,30]. With this number of components, the MCR-ALS analysis was performed through the MATLAB® toolbox developed by Jaumot et al. [31]. Initial guesses for MCR-ALS were spectral type, obtained via the toolbox's built-in PURE initial estimation method [31]. MCR-ALS analyses encompassed the full recorded energy range of the XAS spectra. The principle of the MCR-ALS methodology, as well as more detailed information on the MCR-ALS analyses are provided in the Supporting Information section (Figs. S1 and S2, and related text).

2.3.2. ^{57}Fe Mössbauer spectroscopy

Transmission ^{57}Fe Mössbauer spectra (TMS) of $8\text{Ni}5\text{Fe}/\text{MgAl}_2\text{O}_4$ were collected for the as prepared state, after H_2 -TPR (10 °C/min, up to 800 °C, 30 min) and after DRM (750 °C, 1 atm, 30 min) for $R_c = 1/2$ and $1/1$ at room temperature (RT; 20 °C). For the as prepared state, additional TMS were recorded at -193 °C to obtain better data quality. For surface-sensitive measurements, integrated low-energy electron Mössbauer spectra [32] (ILEEMS) were collected at RT for the as prepared, reduced and spent samples. These ILEEMS were recorded with sample and electron detector (channeltron) contained in a high-vacuum chamber. A bias voltage of + 146 V was applied to the channeltron with respect to the sample holder.

All spectra were recorded in 1024 channels with a velocity scale of approximately ± 10 mm/s. The spectrometer operated in constant acceleration mode with a triangular reference signal and had excellent linearity. A $^{57}\text{Co}(\text{Rh})$ source was used, but, center shift values δ reported hereafter are referenced with respect to $\alpha\text{-Fe}$ at RT. The TMS velocity calibration was based on a RT spectrum of standard hematite ($\alpha\text{-Fe}_2\text{O}_3$), while for ILEEMS it was based on the RT spectrum of an enriched ^{57}Fe foil. For both types of Mössbauer spectroscopy, the velocity (v) increment per channel was 0.0456 ± 0.0001 mm/s. The full line width at half maximum of the inner lines of the calibration spectrum was 0.254 ± 0.002 mm/s. Accumulation of TMS and ILEEMS data took several days until a background of at least 10^6 and 10^5 counts per channel was reached, respectively.

The Mössbauer spectra were fitted with appropriate combinations of Lorentzian profiles representing quadrupole and sextet components, and spectral parameters of these components (center shift (δ), quadrupole splitting (ΔE_Q), full line width at half maximum (Γ), quadrupole shift ($2\varepsilon_Q$) and magnetic hyperfine field (H_{hf})) were determined. These yield information regarding the interactions between the ^{57}Fe nuclei and their electronic and magnetic environment. In turn, this was used to identify

the related Fe phases.

3. Results and discussion

3.1. Speciation of the as prepared state

3.1.1. XAS spectra – as prepared

The XANES spectra at the Fe K and Ni K edge of the as prepared catalyst are presented in Fig. 1. Comparison of the XANES features of the catalyst's initial Fe species with those of reference spectra of $\gamma\text{-Fe}_2\text{O}_3$, $\text{MgFe}^{3+}\text{AlO}_4$ and NiFe_2O_4 (Fig. 1A) reveals an overall Fe^{3+} oxidation state in the sample, as indicated by the similar edge and white line (WL) position. In addition, all Fe K spectra display a pre-edge peak, which is indicative of forbidden quadrupolar $1s \rightarrow 3d$ transitions and d-p orbital mixing associated with dipolar transitions [33]. While all Fe^{3+} reference spectra bear resemblance with the catalyst's as prepared state regarding the shape of the pre-edge and WL features, differences are observed in their intensity. An LCF analysis (Fig. S3) estimates contributions of $\sim 48\%$ NiFe_2O_4 and $\sim 52\%$ $\text{MgFe}^{3+}\text{AlO}_4$, while $\gamma\text{-Fe}_2\text{O}_3$ is not resolved.

The as prepared state's Ni K edge position indicates an overall Ni^{2+} oxidation state (Fig. 1B). The pre-edge features for the Ni species largely overlap with those of NiO in terms of shape, position and intensity. However, the catalyst's WL position is found ~ 0.7 eV higher than that of said NiO reference. While the WL position is closer to that of NiFe_2O_4 (~ 0.3 eV lower) than that of NiAl_2O_4 (~ 0.9 eV lower), the catalyst's pre-edge feature has little resemblance with the former.

Though it is known that discrepancies between reference and catalytic samples can arise from inherent differences in electronic and structural properties between the 'bulk' powder references and the actual catalyst, this mostly has limited effect on the XANES features. It is thus more likely that the observations in Fe K spectra are the result of a combination of spectral contributions of Fe phases. As such, discrepancies observed in the Ni K spectra could be ascribed to the presence of Ni species in NiO, apart from contributions of NiFe_2O_4 [34] and/or NiAl_2O_4 [35]. In accordance, LCF of the as prepared state (Fig. S4) indicates contributions of $\sim 78\%$ NiO, with the remaining $\sim 22\%$ belonging to NiFe_2O_4 .

3.1.2. Mössbauer spectroscopy – as prepared

To further specify the nature of Fe in the as prepared catalyst, TMS and ILEEMS were performed. At RT (Fig. S5), a poorly developed magnetic six-line component superimposed on a doublet is noticed. Fitting with a combination of a model-independent quadrupole-splitting distribution (QSD) and a magnetic hyperfine-field distribution (HFD) yields a relative area (RA) ~ 0.33 for the contribution of the magnetic component to the total spectrum. The hyperfine parameters of both the sextet and doublet component are ill-defined and not conclusive as to the nature of the material.

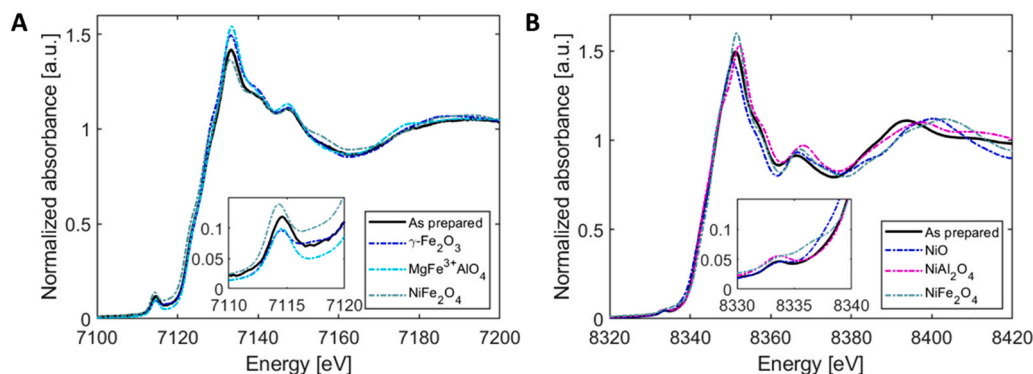


Fig. 1. XANES spectra of as prepared $8\text{Ni}5\text{Fe}/\text{MgAl}_2\text{O}_4$ and selected oxide references at (A) the Fe K edge and (B) the Ni K edge. Reference spectra are shown for comparative purposes. The insets show zooms of the pre-edge features.

At $-193\text{ }^{\circ}\text{C}$ (Fig. 2A), the sextet component is more pronounced and a similar fit using a QSD and a HFD yields an RA value of ~ 0.41 for the magnetic contribution, which is not drastically higher than the RA value at RT. The involved magnetic phase likely possesses a broad transition-temperature range close to RT, implying that at RT the magnetic hyperfine field exhibits a wide distribution and that a fraction of that phase may even be (super)paramagnetic and contribute to the doublet component [36].

At cryogenic temperature, the average hyperfine field ($H_{\text{hf,av}}$) of the magnetic phase is 473 kOe and the maximum-probability field ($H_{\text{hf,m}}$) 486 kOe. There is no obvious fine structure in the calculated distribution profile for the hyperfine field ($p(H_{\text{hf}})$). The average isomer shift is found to be 0.38 mm/s and the quadrupole shift is zero within experimental error limits. These data are consistent with a magnetic phase due to an (Fe,Ni)-rich spinel phase [37], presumably NiFe_2O_4 .

The doublet component in the low-temperature spectrum is adequately described by a QSD with lower and upper limit for ΔE_Q of 0.20 and 1.60 mm/s, respectively. The average isomer shift amounts to $\delta = 0.39\text{ mm/s}$, suggesting trivalent Fe in mixed octahedral O_6 and tetrahedral O_4 coordination. From the calculated probability distribution profile, $p(\Delta E_Q)$ (Fig. S6), an average value for ΔE_Q of $\sim 0.90\text{ mm/s}$ is found. However, the distribution is clearly bimodal, confirming the presence of Fe^{3+} at two different crystallographic sites of the involved Fe-containing phase. This non-magnetic phase can be ascribed to an Fe^{3+} -substituted MgAl_2O_4 spinel [38], proving that impregnation of Fe onto MgAl_2O_4 can lead to its partial incorporation upon calcination. Fe then undergoes interaction with the support, creating spinels of the type $\text{MgFe}^{3+}_x\text{Al}_{2-x}\text{O}_4$, where x varies between 0 and 1. Prior studies [23] have only reported Fe^{3+} in octahedral positions, whereas the current sample has mixed octahedral and tetrahedral coordination. This can relate to the incipient wetness impregnation of Fe and Ni on the MgAl_2O_4 support used in the preparation of this sample, which is in contrast to other works, wherein the Fe, Mg and Al precursors were used in a one-pot coprecipitation synthesis. Notable in the present results is the lack of a sextet component associated with Fe_2O_3 . Since the hyperfine Mössbauer parameters of Fe^{3+} in weakly ferromagnetic spin states related to Fe_2O_3 [39] differ from those associated with the aforementioned spinel phase, Mössbauer spectroscopy has hereby resolved the overlap issue between $\gamma\text{-Fe}_2\text{O}_3$ and NiFe_2O_4 in favor of the last, in line with the LCF analysis of the Fe K edge XANES data.

Summarizing, TMS indicates – in good approximation to LCF analysis of the Fe K edge XANES spectrum – that $\sim 33\text{--}41\%$ of Fe in the as prepared sample is present as a NiFe_2O_4 spinel phase, which orders magnetically, and $\sim 59\text{--}67\%$ as an Fe^{3+} -substituted MgAl_2O_4 spinel phase. Based on prior results [22], this latter phase will hereafter be

referred to as $\text{MgFe}^{3+}_x\text{Al}_{2-x}\text{O}_4$.

To evaluate potential differences between surface and bulk properties of Fe in the catalyst's as prepared state, ILEEMS was performed (Fig. 2B). The spectrum shows a weak emission of only ca. 2%, most likely with a doublet fine structure. Due to the poor statistics, the presence of a magnetic phase could not be verified and the spectrum was analyzed using one single doublet, yielding parameter values as listed in Table 1. The center shift of 0.33 mm/s indicates the valence state of Fe to be Fe^{3+} . Taking into account the center shift and quadrupole splitting values and considering the broad line width, the doublet can actually be composed of two major components arising from Fe^{3+} that substitutes in the octahedral and tetrahedral sites in the MgAl_2O_4 spinel phase. While NiFe_2O_4 is reported as a bulk phase, an associated sextet component cannot be distinguished in ILEEMS.

In summary, Mössbauer spectroscopy indicates an Fe oxidation state of Fe^{3+} in the catalyst's as prepared state, corroborating XAS results. As Mössbauer spectroscopy identifies only $\text{MgFe}_x\text{Al}_{2-x}\text{O}_4$ and NiFe_2O_4 , but does not discern Fe_2O_3 , this relieves the ambiguity of XRD [20]. More so, the confirmed presence of NiFe_2O_4 can account for the observed discrepancies between the as prepared catalyst's XAS spectra and the references in Fig. 1. Further, this identifies Ni present as NiFe_2O_4 , at least partially, as the molar Ni/Fe ratio in $8\text{Ni}5\text{Fe}/\text{MgAl}_2\text{O}_4$ exceeds 1/2. Hence, the additional presence of NiO and/or NiAl_2O_4 must be considered. A schematic representation of the as prepared catalyst state is given in Fig. 3.

3.2. Speciation during and after H_2 -TPR

3.2.1. In situ QXAS

Fig. 4 reports the XANES region of the QXAS spectra recorded during H_2 -TPR at the Fe K and Ni K edge. Reduction takes place for both Fe and Ni atoms, as evidenced by the decrease in WL intensity, an edge shift to lower energies and an increase of the pre-edge peak intensity.

Inspection of the catalyst's final Fe K and Ni K edge spectra reveals that these do not fully correspond with the associated metallic reference spectra. In part, this can be attributed to the final high temperature,

Table 1

Mössbauer parameters of the Fe^{3+} doublet (D1) fitted to the ILEEMS recorded at RT of as prepared $8\text{Ni}5\text{Fe}/\text{MgAl}_2\text{O}_4$ ^a.

Component	δ (mm s ⁻¹)	ΔE_Q (mm s ⁻¹)	Γ (mm s ⁻¹)
D1	Fe^{3+}	0.33(2)	0.89(4)

^a δ : center shift (relative to $\alpha\text{-Fe}$ at $20\text{ }^{\circ}\text{C}$); ΔE_Q : quadrupole splitting; Γ : full line width at half maximum.

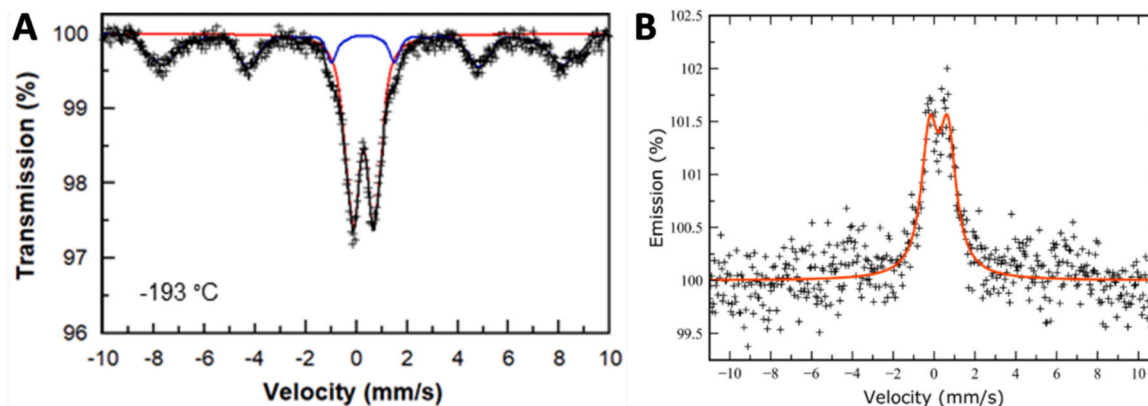


Fig. 2. A: Transmission Mössbauer spectra of as prepared $8\text{Ni}5\text{Fe}/\text{MgAl}_2\text{O}_4$ recorded at $-193\text{ }^{\circ}\text{C}$. + : experimental data; black solid line: spectrum fit; blue solid line: S1 sextet sub-spectrum (Ni-Fe)-rich spinel phase); red solid line: D1 doublet sub-spectrum (Fe^{3+} -substituted MgAl_2O_4 spinel in octahedral and tetrahedral positions). B: ILEEMS of as prepared $8\text{Ni}5\text{Fe}/\text{MgAl}_2\text{O}_4$ at room temperature. + : experimental data; red solid line: spectrum fit, coinciding with the D1 doublet spectrum (Fe^{3+} -substituted MgAl_2O_4 spinel in octahedral and tetrahedral positions).

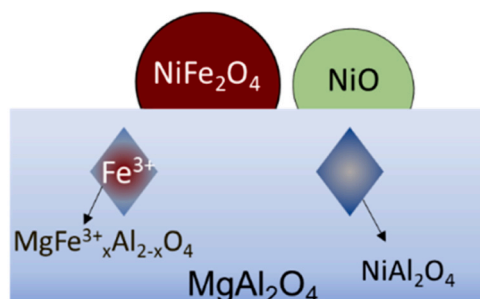


Fig. 3. Schematic representation of the Fe components, as confirmed by Mössbauer spectroscopy, and Ni components, as suggested by XANES, in as prepared 8Ni5Fe/MgAl₂O₄.

800 °C, which entails damping of the features beyond the edge, and to the fact that Ni and Fe species are present within the sample as supported nanoparticles [20,22,40,41], which differ in properties from the ‘bulk’ powder reference species. Furthermore, while it is evident that reduction does occur at both edges, it is plausible that it remains incomplete. Prior XRD measurements in reduced state [20] evidenced the formation of a Ni-Fe alloy phase, next to the support phase. On the other hand, an Fe-modified support MgFe_xAl_{2-x}O₄ such as identified by Mössbauer spectroscopy, was found to reduce partially by H₂-TPR [22, 23]. At the Ni K edge, it is likely that there is still some residual oxidized Ni at the end of reduction. As the metals are supported on an oxide, the presence of a Ni^{δ+} phase, which ensures strong metal-support interactions [22,42] (SMSI), is plausible. SMSI have been reported for different MgAl₂O₄-supported Ni catalysts [43–46], along with Ni-Fe systems [22]. In these cases, SMSI were hypothesized to form a Ni-O bond between the Ni-Fe alloy particle and the support, which remained present after H₂-TPR. On the other hand, the presence of a NiAl₂O₄ phase, remaining stable under the imposed TPR conditions, cannot be excluded either [20].

To differentiate the components in the catalyst during H₂-TPR, PCA and MCR-ALS were applied to the in situ XAS datasets. Applying PCA to the Fe K edge H₂-TPR dataset detects 3 principal components (PCs) (Fig. S7), while 2 PCs are detected for the Ni K edge (Fig. S8). The component spectra and concentration profiles derived by the MCR-ALS methodology are presented in Fig. 5. Associated goodness of fit parameters are provided in Table S1.

On the Fe K edge dataset (Fig. 5A), MCR-ALS allows extracting spectra of 3 different components, with a starting component which decreases in concentration at ~280 °C, a final component which starts to form at a similar temperature and an intermediate which is maximum at about 620 °C with ca. 40% concentration. Considering the known

initial (from Mössbauer spectroscopy) and final [20,22,40] states of Fe within the sample, 4 different components are at play during H₂-TPR: NiFe₂O₄, MgFe³⁺_xAl_{2-x}O₄, its reduced counterpart MgFe²⁺_xAl_{2-x}O₄ [22] together with Ni-Fe alloy [22,40,41,47]. However, the MCR-ALS profile detects only one component at the start of reduction. This is caused by the fact that MCR-ALS can only distinguish species that evolve via different routes, yielding a different variance profile [48]. As a consequence, the Fe component derived by MCR-ALS is actually a ‘merged’ version of multiple Fe phases with different physicochemical nature, in this case NiFe₂O₄ and MgFe³⁺_xAl_{2-x}O₄, which evolve similarly during TPR [24,49]. By MCR-ALS, they will be “merged” into one combi-phase with a spectrum corresponding to a mixture of the two. A one-on-one identification of the derived Fe K edge component spectra is therefore not possible from XAS data alone. At the end of TPR, still under H₂ at 800 °C, a single component appears to be present, with a XANES spectrum resembling the one of metal Fe, which might hence be linked to the Ni-Fe alloy. As for the intermediate state, appearing from 280 °C onwards, this could well originate from Fe²⁺ species, both in the support MgFe_xAl_{2-x}O₄ and as intermediate in partially reduced NiFe₂O₄.

For the Ni K edge dataset, only two components are deduced (Fig. 5B), corresponding to the initial and final state of the catalyst in the experiment. Herein, a slow change in the compositional fractions precedes a noticeable steep increase/decrease starting at ~500 °C. Given the fact that the former change is rather small in comparison to the latter and the fact that the slow rise occurs from the start of the TPR profile onwards, it is probable that these are the result of a temperature effect rather than a change in chemical nature, i.e. reduction, of the Ni species. Above 500 °C, this temperature effect is still present, yet is now dominated by actual reduction of Ni species. A similar observation, though less obvious, can be made for the Fe species, where a weak slope is noticed in the concentration profiles for the Fe components below 280 °C. In view of the as prepared state’s characterization results, component 1 corresponds to a superposition of Ni species in NiFe₂O₄ and NiO. For component 2, metallic Ni is likely the dominant contribution. An SMSI-related NiO phase contribution could persist but possibly in too small a concentration to be detectable. Additionally, as this method relies on variation in the data, it cannot be excluded that a NiAl₂O₄ phase, invariant below 900 °C [50], is equally present but goes unnoticed by MCR-ALS in this reduction process. Numerical experiments (Fig. S9) confirmed that the presence of a small amount (< 10%) of NiAl₂O₄ indeed cannot be excluded.

3.2.2. Mössbauer spectroscopy – reduced

To further assess the identity of Fe species in the reduced sample, ex situ TMS were recorded at RT after reduction, as shown in Fig. 6A. The spectrum is adequately described by a superposition of two Lorentzian-

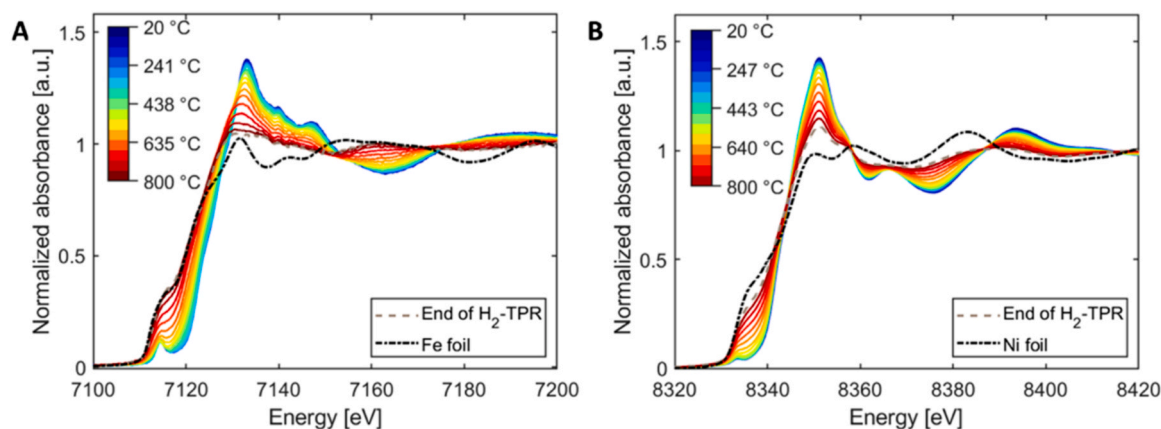


Fig. 4. In situ QXANES spectra recorded during H₂-TPR (10 °C/min, up to 800 °C, for 30 min) of as prepared 8Ni5Fe/MgAl₂O₄ at (A) the Fe K edge and (B) the Ni K edge.

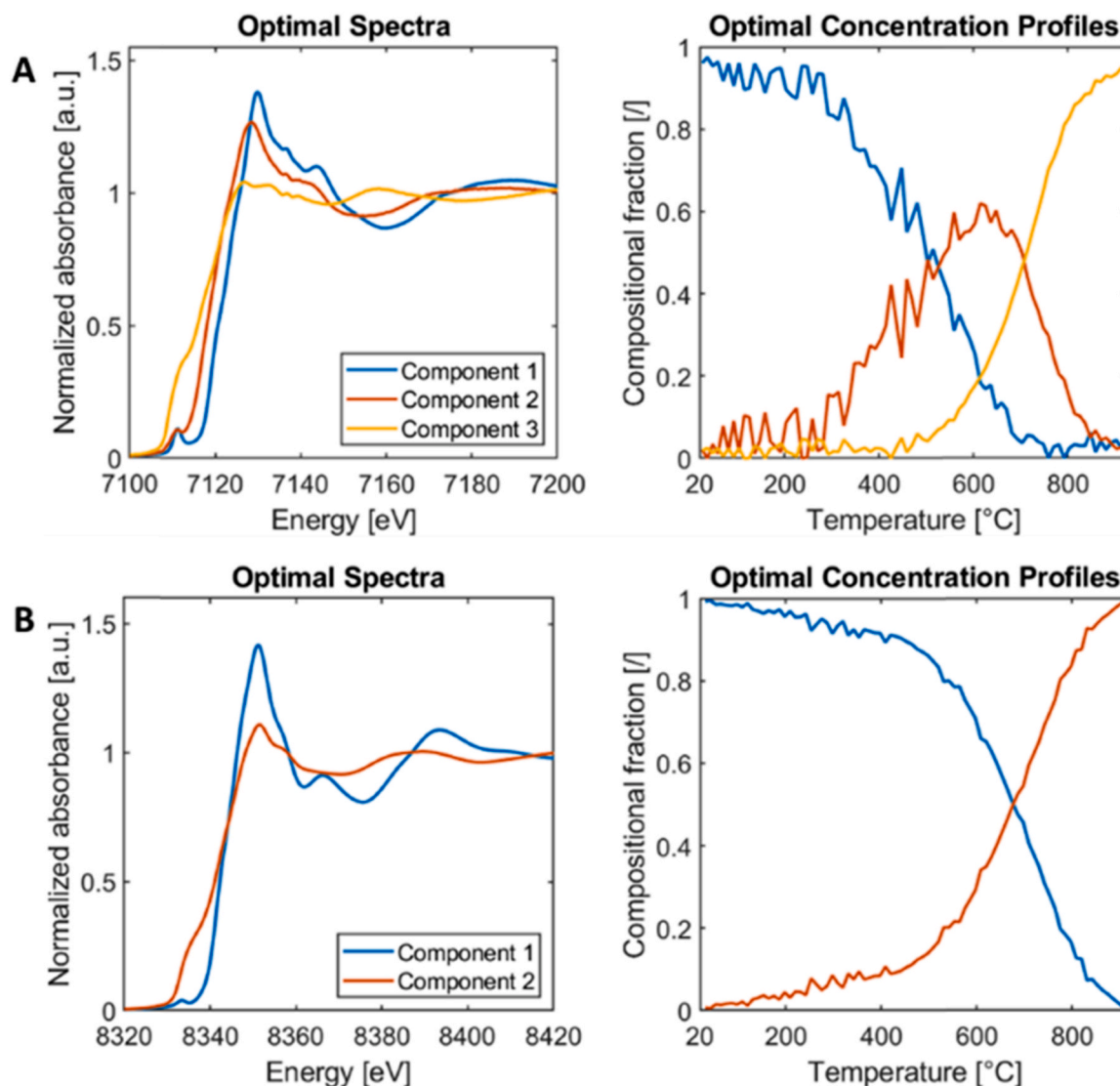


Fig. 5. MCR-ALS results for in situ QXAS data recorded at (A) the Fe K edge and (B) Ni K edge during H_2 -TPR (10 °C/min, up to 800 °C, for 30 min) of as prepared $8Ni5Fe/MgAl_2O_4$. The component spectra are limited to the XANES region for ease of interpretation.

shaped sextets (S1 and S2), an Fe^{3+} quadrupole doublet (D1) and an Fe^{2+} quadrupole doublet (D2). The adjusted parameter values are listed in Table 2.

The values of the hyperfine parameters for S1 and S2 suggest that both sextets are due to two Ni-Fe alloy phases with different Ni/Fe composition [51]. This is plausible considering the nature of the freshly

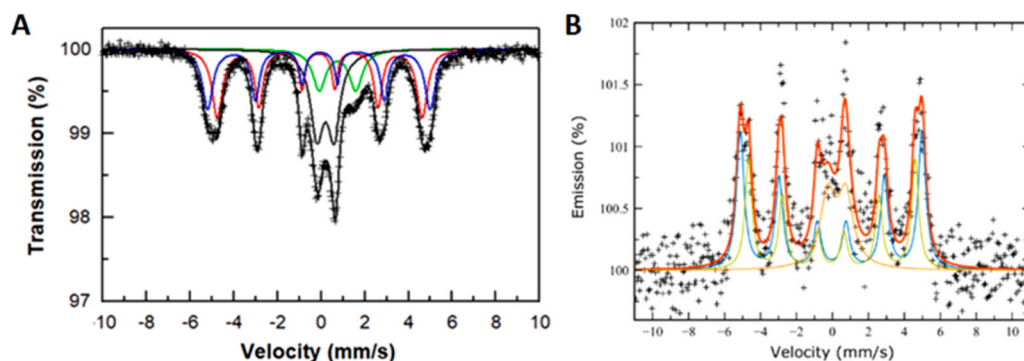


Fig. 6. A: Transmission Mössbauer spectra of reduced $8Ni5Fe/MgAl_2O_4$ recorded at RT. + : experimental data; black solid line: spectrum fit; red solid line: S1 sextet sub-spectrum (Ni-Fe alloy with low at% Ni); blue solid line: S2 sextet sub-spectrum (Ni-Fe alloy with high at% Ni); black solid line: D1 doublet sub-spectrum (Fe^{3+} -substituted $MgAl_2O_4$ spinel in octahedral and tetrahedral positions); black solid line: D2 doublet sub-spectrum (Fe^{2+} -substituted $MgAl_2O_4$ spinel in octahedral and tetrahedral positions). B: Experimental (+) and calculated (red solid line) ILEEMS of reduced $8Ni5Fe/MgAl_2O_4$ recorded at RT. + : experi-

mental data; red solid line: spectrum fit; blue solid line: S1 sextet sub-spectrum (Ni-Fe alloy with low at% Ni); green solid line: S2 sextet sub-spectrum (Ni-Fe alloy with high at% Ni); orange solid line: D1 doublet sub-spectrum (Fe^{3+} -substituted $MgAl_2O_4$ spinel in octahedral and tetrahedral positions).

Table 2

Mössbauer parameters of the two sextets (S1 and S2), the Fe³⁺ doublet (D1) and Fe²⁺ doublet (D2) fitted to the TMS of reduced 8Ni5Fe/MgAl₂O₄ at RT.^a

Component		H_{hf} (kOe)	$2\epsilon_Q$ (mm s ⁻¹)	δ (mm s ⁻¹)	ΔE_Q (mm s ⁻¹)	RA (%)
S1	Fe ⁰	315	-0.05	0.04		29
S2	Fe ⁰	291	0.06	0.02		33
D1	Fe ³⁺			0.31	0.78	25
D2	Fe ²⁺			0.88	1.68	12

^a In case of distributions, the parameters mentioned are those with the highest probability. H_{hf} : magnetic hyperfine field; $2\epsilon_Q$: quadrupole shift; δ : center shift (relative to α -Fe at 20 °C); ΔE_Q : quadrupole splitting; RA: relative area.

prepared parent oxide phases and the broad compositional range of the Ni-Fe alloy as indicated by the Ni-Fe system phase diagram [52]. Johnson et al. [53] reported hyperfine fields gradually varying at RT from ~300 kOe to ~280 kOe upon increasing the at% Ni from 35 to more than 90. According to these results, S1 would arise from a Ni-Fe alloy with relatively low at% Ni, and S2 from a Ni-Fe alloy with higher at% Ni. However, whether it really concerns two distinct alloys is not irrefutable; the two resolved sextets might actually represent a first approximation for the existence of Ni-Fe alloy with a broad range for the Ni/Fe ratio. Attempts to fit the six-line component using a HFD yielded goodness-of-fit values that were significantly higher than for the model using two discrete sextets.

The presence of both an Fe²⁺ and Fe³⁺ quadrupole doublet in the reduced sample, linked to the Fe-modified support MgFe_xAl_{2-x}O₄, is in line with previous findings from QXAS measurements [22]. The line widths for the fitted ferrous and ferric doublets are quite broad, i.e., $\Gamma \approx 0.75$ mm/s, compared to ~0.27 mm/s for standard α -Fe at RT, suggesting that both resolved doublets are each composed of at least two major contributions. Likely, these two would arise from octahedral and tetrahedral Fe²⁺ and Fe³⁺, respectively, in the parent spinel. Note that the adjusted values of δ and ΔE_Q are in line with those obtained for the doublet found for the freshly prepared oxide.

The ILEEMS of the reduced catalyst is reproduced in Fig. 6B. Again, a weak emission of ca. 1.5% is detected. However, the appearance of a six line pattern is obvious, indicating the presence of some magnetic ordering in the particles' surface layers. The spectrum was analyzed using two sextets (S1 and S2, both with imposed line area ratios of 3:2:1 for outer to middle to inner emission lines) and one symmetric Fe³⁺ quadrupole doublet (D). The adjusted Mössbauer parameters are given in Table 3. The sextet parameters are close to those obtained from TMS using a similar fitting model, indicative of Ni-Fe alloys with a broad range for the Ni/Fe ratio at the reduced sample's surface [53]. Taking into account the poor statistics, the doublet parameters probably suggest that the resolved doublet, like for the as prepared sample, is also composed of at least two major components, possibly arising from octahedral and tetrahedral Fe³⁺ in the spinel phase. While an

Table 3

Mössbauer parameters of the two sextets (S1 and S2) and the Fe³⁺ doublet (D1) fitted to the ILEEMS of reduced 8Ni5Fe/MgAl₂O₄ at RT.^a

Component		H_{hf} (kOe)	$2\epsilon_Q$ (mm s ⁻¹)	δ mm (s ⁻¹)	ΔE_Q (mm s ⁻¹)	Γ (mm s ⁻¹)	RA (%)
S1	Fe ⁰	313(1)	-0.03 (1)	0.053 (9)		0.51 (5)	45
S2	Fe ⁰	287(1)	0.06(3)	0.04 (1)		0.47 (4)	32
D	Fe ³⁺			0.35 (4)	0.99(6)	1.0(1)	23

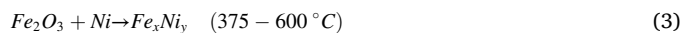
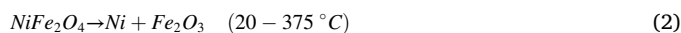
^a In case of distributions, the parameters mentioned are those with the highest probability. H_{hf} : magnetic hyperfine field; $2\epsilon_Q$: quadrupole shift; δ : center shift (relative to α -Fe at RT); ΔE_Q : quadrupole splitting; Γ : full line width at half maximum; RA: relative area.

Fe²⁺-substituted MgAl₂O₄ spinel is not resolved, it is possible that, given the small RA value in TMS, an associated contribution in ILEEMS is overwhelmed by noise.

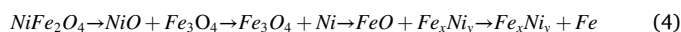
It can be concluded that both the surface and bulk Fe species of the reduced sample comprise similar components: a Ni-Fe alloy with a broad range for the Ni/Fe ratio and the Fe-substituted MgFe_xAl_{2-x}O₄ spinel phase, with both Fe³⁺ and Fe²⁺ presence. Comparison of the RA values of TMS and ILEEMS reveals that the contributions of the alloys are higher for ILEEMS than for TMS. This can be explained through the fact that Ni-Fe alloys are mainly created through the reduction of surface NiFe₂O₄ species by exposure to the reducing environment. As such, the surface is enriched in alloy phase, which leads to relatively higher RA values when analyzing the sample with surface-sensitive techniques. Additionally, the co-existence of Fe²⁺- and Fe³⁺-substituted MgAl₂O₄ support phases is discerned in the reduced catalyst. This is in line with the partial reduction of MgFe³⁺_xAl_{2-x}O₄ into MgFe²⁺_xAl_{2-x}O₄ [22–24].

3.2.3. Discussion

The TMS and ILEEMS results of both the as prepared and reduced samples suggest the presence of different Fe species throughout H₂-TPR of an as prepared 8Ni5Fe/MgAl₂O₄ catalyst. At the start, NiFe₂O₄ and MgFe³⁺_xAl_{2-x}O₄ are present, while MgFe²⁺_xAl_{2-x}O₄, and Ni-Fe alloy (with a dual distribution of high and low Ni/Fe ratios) are discerned after completion of the reduction experiment. From its absence in XRD [20] and Mössbauer spectra in the reduced state, it is evident that NiFe₂O₄ reduces completely into a Ni-Fe alloy. However, reported reduction mechanisms for the aforementioned spinel vary with its preparation method. Benrabaa et al. [54] examined NiFe₂O₄ prepared by two methods: co-precipitation and hydrothermal synthesis. When subjected to H₂-TPR, both types of NiFe₂O₄ displayed a reduction mechanism consisting of consecutive reduction of Ni and Fe:



Chamoumi and co-workers [55] investigated samples synthesized by solid-state reaction. A mechanism was proposed wherein NiFe₂O₄ is first reduced to Ni and FeO at ~460 °C, followed by a final reduction step of FeO to metallic Fe, which then forms a Ni-Fe alloy upon interaction with metallic Ni. In the work of Zhang et al. [56], the decomposition of NiFe₂O₄ into NiO and Fe₃O₄ is reported, followed by a subsequent reduction of these subcomponents. The reduction mechanism they proposed in the range 300 – 550 °C was of the form (with increasing temperature):



Investigations of NiFe₂O₄ supported on CeZrO₂ were performed by Dharanipragada et al. [57]. Herein, in situ XRD during H₂-TPR shows no NiO or Fe oxide, suggesting that NiFe₂O₄ is directly reduced to metallic Fe and Ni in the temperature range 200 – 700 °C. Ma et al. [49] also investigated NiFe₂O₄ supported on CeZrO₂, ZrO₂ and CeO₂, and concluded that all samples reduce similarly in the temperature range of 200 – 900 °C. In contrast to Dharanipragada et al. [57], they proposed a mechanism of the form NiFe₂O₄ → Ni + Fe₃O₄ → NiFe.

The present in situ QXAS data indicate an onset of reduction at temperatures that are lower for Fe species (~280 °C) than for Ni species (~500 °C). Considering the MCR-ALS results at both edges, no formal identification of any of the abovementioned reduction mechanisms is possible. While it is conceivable that the second component derived from the Fe K edge data stems from an intermediate - as implied by its concentration profile, e.g. Fe₃O₄ or FeO -, the merging of components with equal evolution prevents a one-on-one physicochemical identification. In addition and concomitant to NiFe₂O₄ reduction [49], partial reduction of MgFe³⁺_xAl_{2-x}O₄ to MgFe²⁺_xAl_{2-x}O₄ takes place [22–24], as evidenced by TMS. However, the relative presence of Fe in MgFe³⁺_xAl_{2-x}O₄ changes upon reduction: from 59% to 67% for the as

prepared catalyst to roughly 37% for the combination of $\text{MgFe}^{3+}_x\text{Al}_{2-x}\text{O}_4$ and $\text{MgFe}^{2+}_x\text{Al}_{2-x}\text{O}_4$ according to the TMS analysis. Indeed, it has been reported [222] that H_2 -TPR can induce hydrogen spillover from the reduced catalyst's surface species onto the support, thus enabling the migration of support Fe to the surface alloy phase [222]. The migration of Fe from $\text{MgFe}_x\text{Al}_{2-x}\text{O}_4$ during reduction has also been confirmed by Longo et al. [58]. On the basis of Rietveld refinement, they found that $\sim 12\%$ of MgFeAlO_4 segregated as MgFeO_x , while the remaining part of the MgFeAlO_4 spinel phase was retained, albeit with partial reduction of Fe^{3+} to Fe^{2+} . It is thus possible that this MgFeO_x phase also forms during reduction of $8\text{Ni}5\text{Fe}/\text{MgAl}_2\text{O}_4$, as a precursor for Fe migrating out of the support.

The partially reduced support goes unnoticed in the MCR-ALS analysis of the in situ QXAS data. The latter could be due to the involved Fe fractions being smaller than the ones from Ni-Fe alloy, roughly 1/3 for the combination of $\text{MgFe}^{3+}_x\text{Al}_{2-x}\text{O}_4$ and $\text{MgFe}^{2+}_x\text{Al}_{2-x}\text{O}_4$ based on the TMS analysis. Then again, since TMS was performed ex situ, it is equally possible that the actual Fe fractions in the support during high temperature reduction are even lower, in favor of the Ni-Fe alloy fraction. Hence, two Fe contributions can be considered for Ni-Fe alloy formation: NiFe_2O_4 reduction as main alloy source and possibly migration of Fe out of $\text{MgFe}_x\text{Al}_{2-x}\text{O}_4$ towards reduced Ni.

For the Ni species, the NiFe_2O_4 phase is accompanied by NiO, which both reduce to a metallic Ni state, incorporated into Ni-Fe alloy. H_2 -TPR studies on $\text{Ni}/\text{MgAl}_2\text{O}_4$ indicate that supported NiO starts to reduce at $450\text{--}500^\circ\text{C}$ [59–61], which is in agreement with the concentration profiles derived by MCR-ALS for the Ni K edge spectra (Fig. 5B). Previously, an in situ XRD study of the present $8\text{Ni}5\text{Fe}/\text{MgAl}_2\text{O}_4$ catalyst found NiO to reduce at $\sim 500^\circ\text{C}$, after which alloying with Fe took place. The fact that NiFe_2O_4 and NiO are not differentiated by MCR-ALS further strengthens the hypothesis that the steep increase in the compositional fraction of the Ni components corresponds to the joined reduction of NiFe_2O_4 -related Ni species and NiO. In addition, a stable NiAl_2O_4 contribution cannot be excluded.

Summarizing, the results indicate a reduction mechanism as represented in Fig. 7. Herein, the reduction of Fe species in NiFe_2O_4 and $\text{MgFe}^{3+}_x\text{Al}_{2-x}\text{O}_4$ can be considered to occur quasi-simultaneously. NiFe_2O_4 reduces completely to Ni-Fe alloy via a mechanism that is as of yet unknown, while $\text{MgFe}^{3+}_x\text{Al}_{2-x}\text{O}_4$ reduces partially to $\text{MgFe}^{2+}_x\text{Al}_{2-x}\text{O}_4$. A related migration of Fe to the catalyst surface, which contributes to the alloy formation, is not excluded. Reduction of Ni in NiFe_2O_4 and NiO from 500°C on leads to the formation of Ni^0 , which is embedded in the Ni-Fe alloy. A final, residual fraction of oxidic Ni in SMSI-related Ni–O bonds is plausible, while a small ($< 10\%$) Ni fraction related to invariant NiAl_2O_4 is equally likely.

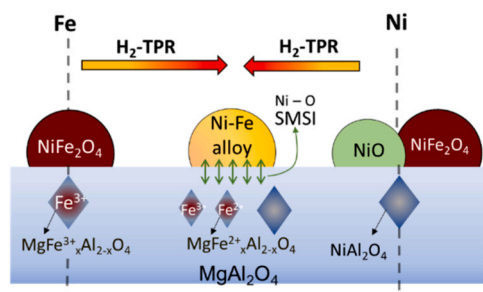


Fig. 7. Schematic representation of the reduction mechanism of as prepared $8\text{Ni}5\text{Fe}/\text{MgAl}_2\text{O}_4$ via H_2 -TPR ($10^\circ\text{C}/\text{min}$, up to 800°C , for 30 min), based on QXAS with MCR-ALS analysis and Mössbauer spectroscopy. Dashed grey lines: as prepared state; \leftrightarrow : SMSI.

3.3. Speciation during and after DRM

3.3.1. In situ QXAS

The current $8\text{Ni}5\text{Fe}/\text{MgAl}_2\text{O}_4$ catalyst achieves in DRM with feed ratio 1/1 a CH_4 conversion of 35% after 4 h TOS and a CO/H_2 ratio of 1.35, as has been reported previously [20]. The present study focuses on linking the behavior of the Ni and Fe species in this catalyst under DRM conditions with this performance. The Fe K edge QXAS spectra recorded during consecutive DRM experiments on a reduced or regenerated catalyst for different CH_4/CO_2 ratios R_c are presented in Fig. 8. The initial Fe K spectra are quasi-identical for all experiments, indicating that post-reaction H_2 reduction can regenerate the catalyst. From the evolution of the edge position and WL intensity, it is clear that oxidation of the Fe species occurs for $R_c < 1/1$. When $R_c = 1/2$, the oxidation appears to be stronger than for $R_c = 1/1.5$, as a higher final WL intensity is observed in the former case, 1.17 vs. 1.15. At $R_c = 1/1$, no net oxidation takes place, although fluctuations in the Fe K edge position occur which are on the order of $\sim 0.3\text{--}0.6\text{ eV}$. For all R_c values, the recorded Ni K edge spectra (Fig. S10) display no net oxidation and are quasi-similar. Yet again, fluctuations appear in the edge position that are on the order of $\sim 0.3\text{--}0.6\text{ eV}$.

PCA analysis of these data discerns 2 PCs at the Fe K edge for $R_c = 1/2$ (Fig. S11) and $1/1.5$ (Fig. S12), while only 1 is derived for $R_c = 1/1$ (Fig. S13). For all corresponding Ni K edge datasets, 1 PC is derived (Figs. S14, S15 and S16), confirming the invariant Ni state. MCR-ALS analysis is thus limited to the $R_c = 1/2$ and $1/1.5$ Fe K data and leads for both datasets to the derivation of identical components (Fig. 9). The corresponding goodness of fit parameters are provided in Table S2.

The first component was identified at the end of reduction as being mainly Ni-Fe, whether or not merged from different alloy compositions, with a possible unresolved contribution from $\text{Fe}^{2+/3+}$ -containing MgAl_2O_4 . During 30 min of DRM, the XAS spectrum for $R_c = 1/2$ and $R_c = 1/1.5$ evolves versus a 60:40 and 90:10 distribution, respectively, between this original reduced component 1 and a component 2, for which a merging effect again cannot be excluded. Based on the WL intensities and pre-edge features of the derived spectral components, it follows that component 2 bears a more oxidized signature. A comparison with reference spectra (Fig. S17) corroborates this, as the component's edge is positioned between those of FeO and Fe_3O_4 . However, its pre-edge feature bears little resemblance to those of oxide references.

Prior in situ XRD experiments for $8\text{Ni}5\text{Fe}/\text{MgAl}_2\text{O}_4$ have elucidated the formation of FeO in DRM under larger CO_2 partial pressures, i.e. $R_c = 1/6$ [20]. For lower values of R_c , this phase was not seen in XRD, which however does not exclude its presence in small concentrations or as crystallites too small to be detected by XRD. XAS, however, is able to detect such oxidized Fe phase even for lower CO_2 concentrations in the feed. This is consistent with the operando XAS results of Kim and co-workers [19], who observed Fe oxide formation during DRM of Ni-Fe/ $\text{Mg}_x\text{Al}_y\text{O}_z$. However, for $R_c = 1/1$, they confirmed FeO formation while the present XAS results do not. This can be related to the MgAl_2O_4 spinel support used in the current study, which is in contrast to the hydrotalcite-derived $\text{Mg}_x\text{Al}_y\text{O}_z$ material employed by Kim et al. [19]. Upon exposure to DRM gases with $R_c < 1/1$, CO_2 indeed induces a net oxidation effect in the Fe species. This oxidation takes place only to a certain extent and within the first $\sim 5\text{--}10\text{ min}$, indicating that the Fe species within the catalyst attain a new steady-state composition under the imposed DRM conditions. Higher steady-state oxide weights are derived for lower R_c values, indicating a more advanced oxidation at higher CO_2 concentrations.

As no noticeable changes are noticed for all the Ni K edge datasets as well as for the $R_c = 1/1$ Fe K edge dataset, it is concluded that these activated catalyst's species don't change significantly under the associated conditions.

3.3.2. Mössbauer spectroscopy – DRM, $R_c = 1/2$

To further identify the Fe species, TMS was performed on an

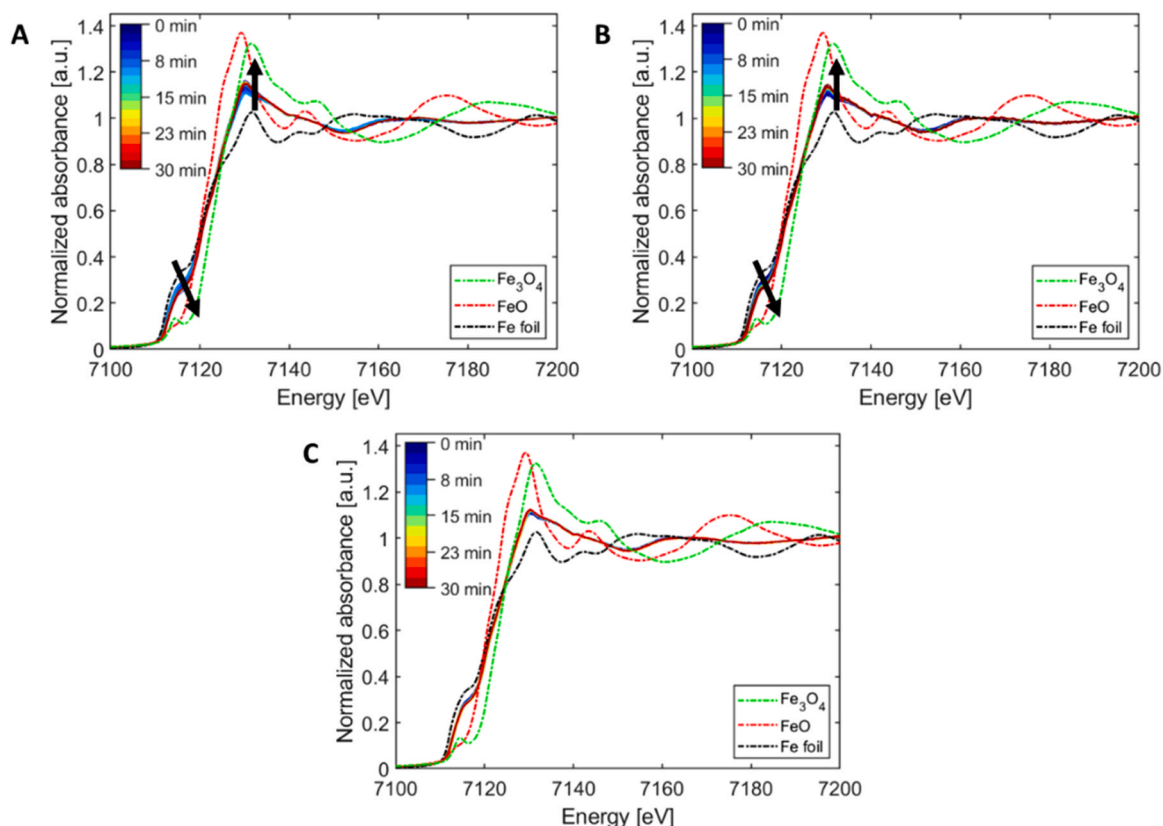


Fig. 8. In situ Fe K edge QXANES spectra recorded during DRM (750 °C, 1 atm, 30 min) of reduced 8Ni5Fe/MgAl₂O₄ with CH₄/CO₂ = (A) 1/2, (B) 1/1.5 and (C) 1/1. Solid lines: experimental spectra; dashed lines: reference spectra; bold arrows: evolution of the spectra during DRM.

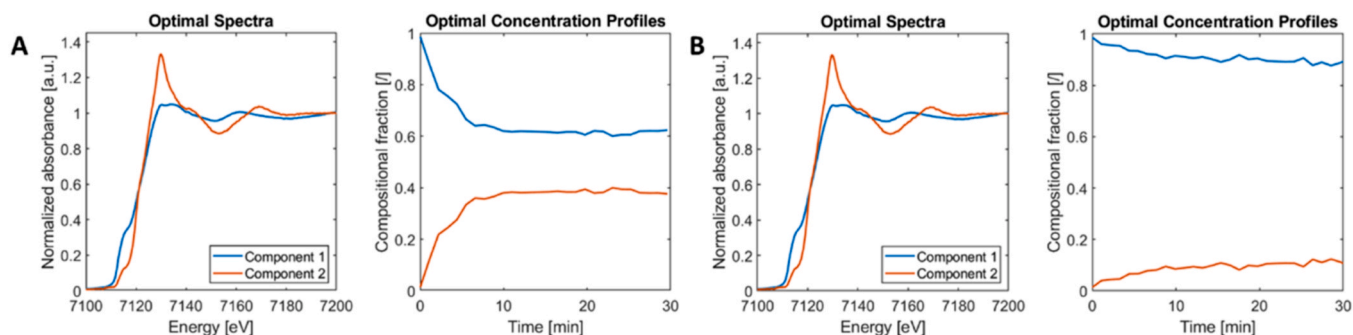


Fig. 9. MCR-ALS results for in situ QXAS data recorded at the Fe K edge during DRM (750 °C, 1 atm, 30 min) with CH₄/CO₂ = (A) 1/2 and (B) 1/1.5 of reduced 8Ni5Fe/MgAl₂O₄. The component spectra are limited to the XANES region for ease of interpretation.

activated sample exposed to DRM conditions with $R_c = 1/2$ (Fig. S18). The spectrum is similar to that recorded for the reduced sample, and is therefore fitted using the same superposition of two sextets (S1 and S2) and two doublets (D1 and D2). The adjusted Mössbauer parameters are collected in Table 4. Compared to TMS for the reduced sample, small differences in terms of RA values are discerned, with a slight increase for the doublet contribution. The latter could point to reincorporation of some Fe into the support lattice. The presence of an additional FeO phase is not resolved, though it is possible that its spectral features are masked in the broadly distributed doublet components [62].

3.3.3. Mössbauer spectroscopy – DRM, $R_c = 1/1$

TMS was also performed for a reduced catalyst exposed to DRM gases with $R_c = 1/1$ to investigate the Fe species under these conditions. The results are provided in Fig. S19. This spectrum is again similar in shape to those of the reduced and DRM-used sample at higher CO₂ partial

Table 4

Mössbauer parameters of the two sextets (S1 and S2), the Fe³⁺ doublet (D1) and Fe²⁺ doublet (D2) fitted to the TMS of 8Ni5Fe/MgAl₂O₄ recorded at RT after DRM (750 °C, 1 atm, 30 min) with CH₄/CO₂ = 1/2.^a

Component		H_{hf} (kOe)	$2\epsilon_Q$ (mm s ⁻¹)	δ (mm s ⁻¹)	ΔE_Q (mm s ⁻¹)	RA (%)
S1	Fe ⁰	315	-0.05	0.03		24
S2	Fe ⁰	289	0.07	0.02		34
D1	Fe ³⁺			0.33	0.75	28
D2	Fe ²⁺			0.83	1.75	14

^a In case of distributions, the parameters mentioned are those with the highest probability. H_{hf} : magnetic hyperfine field; $2\epsilon_Q$: quadrupole shift; δ : center shift (relative to α -Fe at 20 °C); ΔE_Q : quadrupole splitting; RA: relative area.

pressure. Consequently, the spectrum was modelled in the same way and the adjusted parameter values are reported in Table 5.

Within experimental error limits, the hyperfine parameters of the Fe phases in this sample are the same as those of the reduced and the DRM-used sample with non-equimolar gas concentrations. This instigates that the type of phases has not been altered by the prolonged DRM treatment. Nevertheless, marked changes in the RA values are noticeable: these have decreased for both Fe^{2+} and Fe^{3+} doublets in favor of the Ni-Fe alloy sextet components, while keeping the same ratio of Fe^{2+} to Fe^{3+} , i.e. 1:2.

An ILEEMS was also recorded under the same conditions (Fig. 10). As before, the emission is low and the statistics are quite poor. Nevertheless, the appearance of a six line pattern is recognized, indicating that some magnetic ordering occurs in the surface layers of the sample's particles. Contrary to ILEEMS of the reduced sample, only one sextet S could be resolved. In addition, the presence of significant emission centered around zero velocity was apparent and fitted as one symmetric Fe^{3+} quadrupole doublet D.

The relevant adjusted Mössbauer parameters are given in Table 6. Based on the parameter values, it is not feasible to infer correlation between sextets S1 and S2 of the reduced sample and sextet S of this DRM-used sample. Sextet S shows a very broad line width (~ 0.9 mm/s), which is possibly due to Ni-Fe alloys with a broad distribution for the Ni/Fe ratio [53], so this could still match the 2 sextets that were resolved in ILEEMS for the reduced sample (Fig. 6B).

Roughly, the doublet parameters are in accordance with the results obtained for the reduced sample, and the doublet can therefore again be attributed to octahedral and tetrahedral Fe^{3+} in the spinel phase. As before, an Fe^{2+} -substituted MgAl_2O_4 spinel is not resolved. As its RA value in TMS is even smaller than for the reduced sample, 8 vs. 12%, an associated contribution in ILEEMS is not expected to protrude above the noise. Remarkably, the RA values of the doublet are equal for the reduced and the present DRM-used sample, ca. 23%, indicating that the surface composition has not changed after DRM with $R_c = 1/1$.

While the Ni-Fe alloys formed upon reduction are surface species [15,16,20,40,63–65], the increase of the alloy phase's RA with respect to the Fe-modified spinel after DRM is not reflected in ILEEMS, only in TMS. Given the ILEEMS information depth of ~ 5 nm versus the reported size of 14.3 nm for the alloy particles [20], it is plausible that additional Fe alloying goes unnoticed in ILEEMS. On the other hand, the low data quality of ILEEMS will surely entail higher experimental error, also in the RA values.

3.3.4. Discussion

From the collective analysis of Mössbauer spectroscopy and in situ QXAS, it follows that the compositional changes occurring in the activated catalyst during DRM depend on the applied R_c value. These results can be explained through the following considerations. As found by TMS and MCR-ALS, the activated catalyst consists predominantly of Ni-Fe alloy, along with $\text{MgFe}^{2+}_x\text{Al}_{2-x}\text{O}_4$, $\text{MgFe}^{3+}_x\text{Al}_{2-x}\text{O}_4$, and potentially NiAl_2O_4 and SMSI-related NiO as separate Ni phases. Theofanidis et al. [20] suggest DRM on Ni-Fe/ MgAl_2O_4 takes place on the alloy via a

Table 5

Mössbauer parameters of two sextets (S1, S2), the Fe^{3+} and Fe^{2+} doublet (D1, D2, respectively) fitted to the TMS of 8Ni5Fe/ MgAl_2O_4 recorded at RT after DRM (750 °C, 1 atm, 30 min) with $\text{CH}_4/\text{CO}_2 = 1/1$.^a

Component		H_{hf} (kOe)	$2\epsilon_Q$ (mm s^{-1})	δ (mm s^{-1})	ΔE_Q (mm s^{-1})	RA (%)
S1	Fe^0	315	-0.05	0.03		38
S2	Fe^0	291	0.07	0.03		38
D1	Fe^{3+}			0.31	0.7	16
D2	Fe^{2+}			0.85	1.78	8

^a In case of distributions, the parameters mentioned are those with the highest probability. H_{hf} : magnetic hyperfine field; $2\epsilon_Q$: quadrupole shift; δ : center shift (relative to $\alpha\text{-Fe}$ at 20 °C); ΔE_Q : quadrupole splitting; RA: relative area.

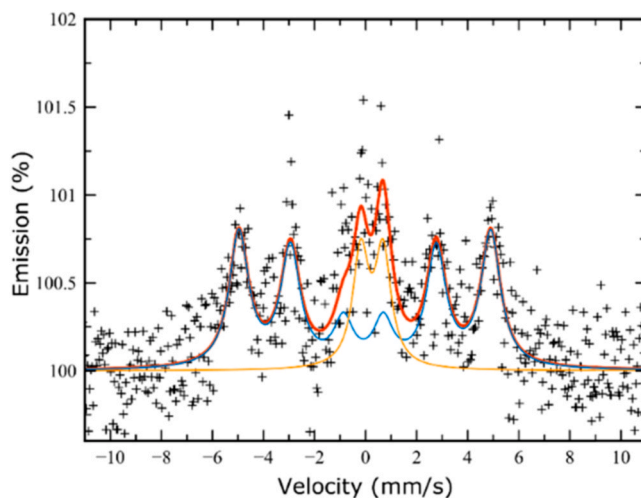


Fig. 10. ILEEMS of 8Ni5Fe/ MgAl_2O_4 after DRM (750 °C, 1 atm, 30 min) with $\text{CH}_4/\text{CO}_2 = 1/1$, recorded at RT. + : experimental data; red solid line: spectrum fit; blue solid line: S sub-spectrum (Ni-Fe alloy with broad Ni/Fe range); orange solid line: D doublet sub-spectrum (Fe^{3+} -substituted MgAl_2O_4 spinel in octahedral and tetrahedral positions).

Table 6

Mössbauer parameters of the sextet (S) and the Fe^{3+} doublet (D) fitted to the ILEEMS of 8Ni5Fe/ MgAl_2O_4 recorded at RT after DRM (750 °C, 1 atm, 30 min) with $\text{CH}_4/\text{CO}_2 = 1/1$.^a

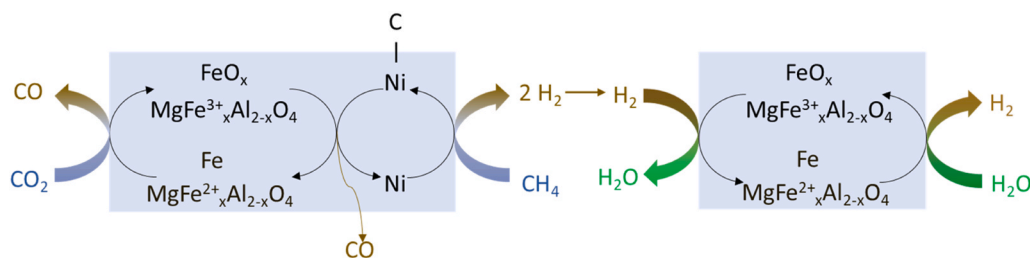
Component		H_{hf} (kOe)	$2\epsilon_Q$ (mm s^{-1})	δ (mm s^{-1})	ΔE_Q (mm s^{-1})	Γ (mm s^{-1})	RA (%)
S	Fe^0	306 (3)	0.06 (5)	0.05 (4)		0.92 (7)	76
D	Fe^{3+}			0.35 (4)	0.87(6)	0.8(1)	24

^a In case of distributions, the parameters mentioned are those with the highest probability. H_{hf} : magnetic hyperfine field; $2\epsilon_Q$: quadrupole shift; δ : center shift (relative to $\alpha\text{-Fe}$ at RT); ΔE_Q : quadrupole splitting; Γ full line width at half maximum; RA: relative area.

Mars-van Krevelen mechanism. Herein, the Ni sites activate the dissociation of CH_4 , while the alloy's constituent Fe sites are oxidized by CO_2 to FeO_x . The latter is subsequently reverted into Fe through reduction by C, CH_x and H species, originating from CH_4 dissociation.

With the confirmed presence of Fe in the MgAl_2O_4 support, this scheme must be extended. Indeed, studies on $\text{Ni}/\text{MgFe}_x\text{Al}_{2-x}\text{O}_4$ have substantiated the redox abilities of $\text{MgFe}^{Y+}_x\text{Al}_{2-x}\text{O}_4$ (where $Y = 2$ or 3) species [22,66]; they can oxidize CH_4 dissociation products and can be oxidized by CO_2 and/or H_2O . On its own though, Fe in $\text{MgFe}^{Y+}_x\text{Al}_{2-x}\text{O}_4$ is far less active in DRM than supported Ni-Fe/ MgAl_2O_4 [20,22]. Hence, a Mars-van Krevelen mechanism is proposed which accounts for the contribution of both this Fe-substituted MgAl_2O_4 and supported Ni-Fe species, as represented schematically in Fig. 11.

Fig. 12 displays the changes during DRM for different values of R_c . In the case of more oxidizing conditions ($R_c < 1$) (Fig. 12B), the CH_4 dissociation products are less abundant, such that a net oxidation of Fe takes place by the more dominant CO_2 . Oxidation of Ni by CO_2 under DRM conditions is unfavorable [20], hence Ni retains its metallic state. According to MCR-ALS of the in situ QXAS data, 40% ($R_c = 1/2$) of Fe gets re-oxidized towards FeO during DRM, leaving 60% in alloyed state. Although the lineshape of component 2 in Fig. 9A confirms an oxidized Fe state, the existence of FeO_x during DRM cannot be confirmed unambiguously with the current results. In comparing with model references, the difference in temperature between the reference (RT) and the catalyst recording (750 °C), and the different nature of the signal



reactants; brown arrow sections: DRM products; green arrow section: H₂O contributions from reverse water-gas shift.

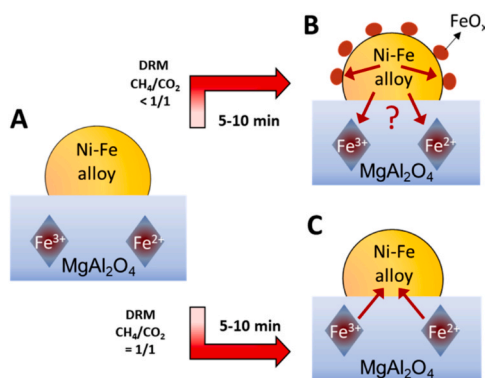


Fig. 12. Schematic representation of the compositional changes of (A) reduced 8Ni5Fe/MgAl₂O₄ during DRM (750 °C, 1 atm, 30 min) with (B) CH₄/CO₂ < 1/1 and (C) CH₄/CO₂ = 1/1, as suggested by in situ QXAS. Fe³⁺ and Fe²⁺ denote MgFe³⁺_xAl_{2-x}O₄ and MgFe²⁺_xAl_{2-x}O₄, respectively. For ease of representation, abstraction is made of NiAl₂O₄ and SMSI-related Ni – O bonds. Red arrows in B and C indicate Fe mobility.

(mixed catalyst and support contributions vs. homogeneous bulk) need to be taken into account. Still, the presence of FeO can be deemed plausible, considering previous studies. Hu et al. [67] found FeO to be present in in situ XRD during DRM with $R_c = 1/2$ for a catalyst with higher Fe loading. For the present loading of 5 wt% Fe, no FeO was discerned by in situ XRD until the R_c was set at 1/6 [20], i.e. a far more oxidizing environment. For higher values of R_c , the XRD-amorphous nature of such FeO particles would explain the absence of the phase in the diffraction patterns [20]. Based on the present QXAS and MCR-ALS, an oxidized phase indeed forms. MCR-ALS indicates this oxidation occurs within the first 5–10 min of the reaction, after which the catalyst attains a steady-state composition for DRM. In addition, this oxidation is deeper for lower R_c values as a consequence of the more oxidative environment.

When looking with ex situ TMS after reaction with $R_c = 1/2$, the DRM effect is limited with the alloy RA being slightly lower than after reduction, 58% vs. 62% respectively. When Fe segregates out of the Ni-Fe alloy to the surface under these DRM conditions, the alloy's bulk gets partially depleted of Fe. Even so, due to the broad Ni/Fe ratio of the alloy phase present, it remains in line with component 1 of the MCR-ALS analysis, the unchanged Ni K edge spectra and the Ni-Fe phase diagram [52]. The concomitant change in doublet RA from TMS, 37–42%, is not very large, although if real, it could indicate that part of the extracted Fe gets reincorporated into the support lattice, like in the calcination process. The presence of FeO on the other hand is not resolved, but still possible given the broad signal. Hence, the oxidized iron phase detected by QXAS and MCR-ALS combines small (< 3 nm) FeO_x nanoparticles at the surface of the Ni-Fe alloy core and possibly some Fe^{2+/3+} reincorporated into the support. Both have proven efficiency in carbon mitigation [20,22].

In a more reducing environment ($R_c = 1/1$) (Fig. 12C), sufficient CH₄

dissociation product species are present to counteract oxidation of Fe species by CO₂, leading to unchanged Fe K edge QXAS data. In line with the higher concentrations of reducing CH₄ relative to the case of $R_c = 1/2$, post-reaction TMS and ILEMS indicate higher contributions of Ni-Fe alloy. Interestingly, TMS post DRM also reveals a higher abundance of Ni-Fe alloy compared to the reduced state. The latter indicates that reduction of the sample was not complete after the H₂-TPR treatment, so that reduction and alloying continued in these DRM reaction conditions, in contrast to reaction with $R_c = 1/2$. This additional alloying under DRM with $R_c = 1/1$ counters Fe extraction from the alloy by CO₂, leading to the very stable performance of 8Ni5Fe/MgAl₂O₄ under such conditions, as established previously [20]. The explanation for the insignificant changes at the Ni K edge is analogous to the case of $R_c < 1$.

On the basis of these results, it is proposed that DRM would benefit from the use of higher CH₄/CO₂ ratios as these would avoid Fe segregation from the Ni-Fe alloy, which would otherwise lead to coverage of active Ni sites.

4. Conclusions

The combined use of in situ QXAS and ⁵⁷Fe Mössbauer spectroscopy has led to novel insights regarding the constituent Ni and Fe phases in 8Ni5Fe/MgAl₂O₄ catalysts during H₂-TPR and DRM.

- The Fe phases in the as prepared catalyst have been identified as NiFe₂O₄ and MgFe³⁺_xAl_{2-x}O₄, while significant Ni contributions have been derived as NiO next to NiFe₂O₄.
- Upon H₂-TPR, NiFe₂O₄ completely reduces, forming a Ni-Fe alloy, to which quasi-complete reduction of NiO contributes. The MgFe³⁺_xAl_{2-x}O₄ phase partially reduces into MgFe²⁺_xAl_{2-x}O₄.
- During DRM, the induced changes are dependent on the applied CH₄/CO₂ ratio and can be explained by a Mars-van Krevelen mechanism.
 - o For CH₄/CO₂ < 1, a net oxidation of Fe occurs, which extracts it from the Ni-Fe alloy, thereby generating FeO_x nanoparticles that decorate the remainder alloy and possibly even reincorporating some Fe into the support lattice.
 - o For a CH₄/CO₂ ratio = 1, CO₂-induced oxidation is compensated by reduction through product H₂ and CO, leading to ongoing alloying.

This interplay between gas environment and active elements/phases, both in alloy and support, is considered responsible for the DRM performance of 8Ni5Fe/MgAl₂O₄.

CRediT authorship contribution statement

Valentijn De Coster: Conceptualization, Methodology, Software, Formal analysis, Writing – original draft, Writing – review & editing. **Nadadur Veeraraghavan Srinath:** Methodology, Formal analysis, Writing – original draft. **Stavros Alexandros Theofanidis:** Investigation, Writing – review & Editing. **Laura Pirro:** Formal analysis, Writing –

original draft. **Antoine Van Alboom**: Mössbauer investigation, Formal analysis, Writing – original draft. **Hilde Poelman**: Conceptualization, Writing – original draft, Writing – review & editing, Supervision, Funding acquisition. **Maarten K. Sabbe**: Supervision. **Guy B. Marin**: Supervision, Funding acquisition. **Vladimir V. Galvita**: Conceptualization, Writing – original draft, Writing – review & editing, Supervision, Funding acquisition.

Declaration of Competing Interest

The authors declare that there are no conflicts of interest.

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Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at [doi:10.1016/j.apcatb.2021.120720](https://doi.org/10.1016/j.apcatb.2021.120720).

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